=> d his

12 S L20 OR L18

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(FILE 'REGISTRY' ENTERED AT 10:46:09 ON 03 JUL 2003)
                DEL HIS Y
                ACT TELLER2/A
        178763) SEA FILE=REGISTRY ABB=ON PLU=ON 333.401/RID STR
SCR 1840
19010 SEA FILE=REGISTRY SUB=L1 SSS FUL L2 NOT L3 Nonoved
L1 (
L2
L3
L4
     FILE 'HCAPLUS' ENTERED AT 10:55:03 ON 03 JUL 2003
L5
          22041 S L4
          71352 S ALBUMIN#
L6
L7
             24 S L5 AND L6
            284 S LIGAND# AND L5
rs
L9
             3 S L8 AND L6
L10
         323170 S ADSOR? OR DESOR?
            152 S L10 AND L5
L11
L12
             2 S L11 AND L6
L13
            783 S ELECTRON? (L) WITHDR?
L14
             3 S L5 AND L13
L15
          11696 S (ELECTRON? (S) WITHDR?)/AB
            53 S L15 AND L5
L16
L17
             2 S L16 AND (L6 OR LIGAND?)
             7 S L9 OR L12 CR L14 OR L17
L18
L19
          7342 S L6 (L) BIND?
L20 _ _ _ 6 S L19 AND L5 _ _
```

L21

=> fil reg FILE 'REGISTRY' ENTERED AT 11:01:38 ON 03 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2003 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 2 JUL 2003 HIGHEST RN 541497-70-5 2 JUL 2003 HIGHEST RN 541497-70-5 DICTIONARY FILE UPDATES:

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

=> d que stat 14

L1 (178763) SEA FILE=REGISTRY ABB=ON PLU=ON 333.401/RID

----2- --- 7- ---

NODE ATTRIBUTES: CONNECT IS E3 RC AT DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS

STEREO ATTRIBUTES: NONE L3 SCR 1840

L419010 SEA FILE=REGISTRY SUB=L1 SSS FUL L2 NOT L3

100.0% PROCESSED 23279 ITERATIONS

SEARCH TIME: 00.00.01

19010 ANSWERS

=> fil hcaplus FILE 'HCAPLUS' ENTERED AT 11:01:51 ON 03 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.

narrowed south of not allowing any

PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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FILE COVERS 1907 - 3 Jul 2003 VOL 139 ISS 1 FILE LAST UPDATED: 2 Jul 2003 (20030702/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

'OBI' IS DEFAULT SEARCH FIELD FOR 'HCAPLUS' FILE

=> d hsi 15-

'HSI' IS NOT A VALID FORMAT FOR FILE 'HCAPLUS'
'L5-' IS NOT A VALID FORMAT FOR FILE 'HCAPLUS'
ENTER DISPLAY FORMAT (BIB):end

=> d his 15-

(FILE 'REGISTRY' ENTERED AT 10:46:09 ON 03 JUL 2003)

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FILE 'HCAPLUS' ENTERED AT 10:55:03 ON 03 JUL 2003
L5
          22041 S L4
L6
          71352 S ALBUMIN#
L7
             24 S L5 AND L6
L8
            284 S LIGAND# AND L5
L9
              3 S L8 AND L6
L10
         323170 S ADSOR? OR DESOR?
            152 S L10 AND L5
L11
L12
              2 S L11 AND L6
            783 S ELECTRON? (L) WITHDR?
L13
L14
              3 S L5 AND L13
          11696 S (ELECTRON? (S) WITHDR?)/AB
L15
             53 S L15 AND L5
L16
              2 S L16 AND (L6 OR LIGAND?)
L17
              7 S L9 OR L12 OR L14 OR L17
L18
           7342 S L6 (L) BIND?
L19
L20
              6 S L19 AND L5
L21
             12 S L20 OR L18
```

FILE 'REGISTRY' ENTERED AT 11:01:38 ON 03 JUL 2003

FILE 'HCAPLUS' ENTERED AT 11:01:51 ON 03 JUL 2003

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=> d que nos 121
L1 ( 178763)SEA FILE=REGISTRY ABB=ON PLU=ON 333.401/RID
L2 STR
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SCR 1840
L3
L4
         19010 SEA FILE=REGISTRY SUB=L1 SSS FUL L2 NOT L3
L5
         22041 SEA FILE=HCAPLUS ABB=ON PLU=ON L4
         71352 SEA FILE=HCAPLUS ABB=ON PLU=ON ALBUMIN#/OBI
L6
           284 SEA FILE=HCAPLUS ABB=ON PLU=ON LIGAND#/OBI AND L5
L8
L9
             3 SEA FILE=HCAPLUS ABB=ON PLU=ON L8 AND L6
        323170 SEA FILE=HCAPLUS ABB=ON PLU=ON ADSOR?/OBI OR DESOR?/OBI
L10
           152 SEA FILE=HCAPLUS ABB=ON PLU=ON L10 AND L5
L11
             2 SEA FILE=HCAPLUS ABB=ON PLU=ON L11 AND L6
L12
L13
           783 SEA FILE=HCAPLUS ABB=ON PLU=ON ELECTRON?/OBI (L) WITHDR?/OBI
L14
             3 SEA FILE=HCAPLUS ABB=ON PLU=ON L5 AND L13
         11696 SEA FILE=HCAPLUS ABB=ON PLU=ON (ELECTRON? (S) WITHDR?)/AB
L15
            53 SEA FILE=HCAPLUS ABB=ON PLU=ON L15 AND L5
L16
             2 SEA FILE=HCAPLUS ABB=ON PLU=ON L16 AND (L6 OR LIGAND?/OBI)
L17
L18
             7 SEA FILE=HCAPLUS ABB=ON PLU=ON L9 OR L12 OR L14 OR L17
L19
          7342 SEA FILE=HCAPLUS ABB=ON PLU=ON L6 (L) BIND?/OBI
             6 SEA FILE=HCAPLUS ABB=ON PLU=ON L19 AND L5
L20
L21
            12 SEA FILE=HCAPLUS ABB=ON PLU=ON L20 OR L18
```

=> d .ca hitstr 121 1-12

L21 ANSWER 1 OF 12 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 2001:396779 HCAPLUS

DOCUMENT NUMBER: 135:10396

A method for anion-exchange adsorption and TITLE:

anion-exchangers

S): Johansson, Bo-lennart; Andersson, Mikael; Gustavsson,

Jan; Belew, Makonnen; Maloisel, Jean-luc INVENTOR(S):

PATENT ASSIGNEE(S): Amersham Pharmacia Biotech Ab, Swed. SOURCE: PCT Int. Appl., 50 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

```
PATENT NO. KIND DATE
                                        APPLICATION NO. DATE
    -----
                                         -----
    WO 2001038227
                    A2
                           20010531
                                        WO 2000-EP11605 20001122
    WO 2001038227
                    A3 20011115
           AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
            HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
            LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
            SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
            YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
            DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
            BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                    A2 20020904
                                       EP 2000-979615 20001122
    EP 1235748
           AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                    T2 20030422
                                         JP 2001-539791
                                                          20001122
    JP 2003514664
                                                    A 19991122
PRIORITY APPLN. INFO.:
                                      SE 1999-4197
                                      WO 2000-EP11605 W 20001122
```

A method for the removal of a substance carrying a neg. charge and being AB

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present in an aq. liq. (I). The method comprises the steps of: (i)
contacting the liq. with a matrix carrying a plurality of ligands
comprising a pos. charged structure and a hydrophobic structure, and (ii)
desorbing the substance. The characterizing feature is that (I) each of
said ligands together with a spacer has the formula: --
SP---[Ar-R1-N+(R2R3R4)] where (A) [Ar-R1-N+(R2R3R4)] represents a ligand
(a) Ar is an arom. ring, (b) R1 is [(L)nR'1]m where n and m are integers
selected amongst zero or 1; L is amino nitrogen, ether oxygen or thioether
sulfur; R'1 is a linker selected among (1) hydrocarbon groups; (2)
-C(=NH)-; (c) R2-4 are selected among hydrogen and alkyls; (B) SP is a
spacer providing a carbon or a heteroatom directly attached to
Ar-R1-N+(R2R3R4); (C) --- represents that SP replaces a hydrogen in
[Ar-R1-N+(R2R3R4)]; (D) -- represents binding to the matrix; and (II)
desorption. There is also described (a) anion-exchangers having high
breakthrough capacities, (b) a screening method and (c) a desalting
protocol.
ICM C02F001-28
ICS B01J041-00
66-4 (Surface Chemistry and Colloids)
Section cross-reference(s): 9, 80
anion exchange adsorption protein recovery sepn; ionic strength
adsorption protein anion exchanger
Allylation
Bromination
Molecular structure-property relationship
   (Sepharose 6 Fast Flow matrix modified by allylation with allyl
   glycidyl ether proceeded by bromination and coupling with various
   nitrogen contg. ligands)
Anion exchange
                 Anion exchangers
Ionic strength
   (Sepharose 6 Fast Flow matrix modified with various ligands
   in which there is a pos. charged nitrogen under conditions permitting
   binding between the anion-exchanger and various proteins at high ionic
   strengths)
Lactalbumins
Proteins, general, properties
RL: ANT (Analyte); PEP (Physical, engineering or chemical process); PRP
(Properties); ANST (Analytical study); PROC (Process)
   (Sepharose 6 Fast Flow matrix modified with various ligands
   in which there is a pos. charged nitrogen under conditions permitting
   binding between the anion-exchanger and various proteins at high ionic
   strengths)
Liquid chromatography
   (Sepharose 6 Fast Flow matrix modified with various ligands
   in which there is a pos. charged nitrogen under conditions permitting
   binding between the anion-exchanger and various proteins at high ionic
  strengths evaluated using)
Adsorption
 Desorption
   (method for anion-exchange adsorption and anion-exchangers
   and desorption from them)
Albumins, properties
RL: ANT (Analyte); PEP (Physical, engineering or chemical process); PRP
(Properties); ANST (Analytical study); PROC (Process)
   (serum; Sepharose 6 Fast Flow matrix modified with various
  ligands in which there is a pos. charged nitrogen under
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conditions permitting binding between the anion-exchanger and

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various proteins at high ionic strengths)
     60-23-1, Cysteamine
                          104-14-3, Octopamine 106-92-3, Allyl glycidyl
TΤ
             3674-06-4
                        6674-22-2, 1,8-Diazabicyclo[5,4,0]-undec-7-ene
     7726-95-6, Bromine, reactions 19406-49-6
                                                67385-09-5
                                                               106894-56-8,
     Fmoc-L-tyrosine-N-hydroxysuccinimide ester
     RL: MOA (Modifier or additive use); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (Sepharose 6 Fast Flow matrix modified by allylation with allyl
        glycidyl ether proceeded by bromination and coupling with various
        nitrogen contg. ligands)
     136109-66-5, sepharose 6 fast flow
TΤ
     RL: AMX (Analytical matrix); PEP (Physical, engineering or chemical
     process); PRP (Properties); ANST (Analytical study); PROC (Process)
        (Sepharose 6 Fast Flow matrix modified with various ligands
        in which there is a pos. charged nitrogen under conditions permitting
        binding between the anion-exchanger and various proteins at high ionic
        strengths)
                             9078-38-0, soybean trypsin inhibitor
IT
     1391-06-6, conalbumin
     RL: ANT (Analyte); PEP (Physical, engineering or chemical process); PRP
     (Properties); ANST (Analytical study); PROC (Process)
        (Sepharose 6 Fast Flow matrix modified with various ligands
        in which there is a pos. charged nitrogen under conditions permitting
        binding between the anion-exchanger and various proteins at high ionic
        strengths)
     127546-40-1, Q Sepharose fast flow
IΤ
     RL: AMX (Analytical matrix); PEP (Physical, engineering or chemical
     process); PRP (Properties); ANST (Analytical study); PROC (Process)
        (Sepharose 6 Fast Flow matrix modified with various ligands
        in which there is a pos. charged nitrogen under conditions permitting
        binding between the anion-exchanger and various proteins at high ionic
        strengths compared with)
IT
     51-41-2, Noradrenaline
                              60-18-4, Tyrosine, reactions
                                                             63-74-1.
                     99-57-0, 2-Amino-4-nitrophenol
     Sulfanilamide
                                                    119-62-0
                                                                 123-30-8.
                     500-88-9, Tyrosinol
                                           526-53-4, Tryptophanol
                                                                    552-85-2
     4-Aminophenol
                                    1004-39-3, 4,6-Diamino-2-
     934-32-7, 2-Aminobenzimidazole
                          1193-02-8, 4-Aminothiophenol
     mercaptopyrimidine
                                                         3204-61-3,
                               3306-06-7, 2-Amino-1-phenyl-1,3-propanediol
     1,2,4,5-Tetraaminobenzene
                13472-00-9, 2-(4-Aminophenyl)ethylamine 16088-07-6
     7621-14-9
     16854-32-3, Thiomicamine
                                36469-86-0 37491-68-2, 3,4-
                          341014-76-4
                                          341014-77-5
     Dihydroxybenzylamine
                                                        341014-78-6
     341032-58-4
     RL: MOA (Modifier or additive use); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (elution cond. for three proteins and breakthrough capacity of BSA on
        Sepharose 6 Fast Flow anion-exchangers modified with ligands
        of)
ΙT
     934-32-7, 2-Aminobenzimidazole
     RL: MOA (Modifier or additive use); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (elution cond. for three proteins and breakthrough capacity of BSA on
        Sepharose 6 Fast Flow anion-exchangers modified with ligands
        of)
     934-32-7 HCAPLUS
RN
```

1H-Benzimidazol-2-amine (9CI) (CA INDEX NAME)

CN

L21 ANSWER 2 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

2000:441826 HCAPLUS

DOCUMENT NUMBER:

133:71091

TITLE:

Removal/purification of serum albumins using

WO 1999-EP10123 W 19991220

matrix-immobilized affinity ligands

INVENTOR(S):

Regberg, Tor; Ellstrom, Christel Amersham Pharmacia Biotech AB, Swed.

PATENT ASSIGNEE(S):

SOURCE:

PCT Int. Appl., 24 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA?	rent	NO.		KI	ND	DATE			A	PPLI	CATI	ON NO	ο.	DATE			
WO	WO 2000037501			A1		20000629		WO 1999-EP10123			19991220						
	W:	AU,	CA,	JP,	US												
	RW:	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,
		PT,	SE														
CA	2355	827		A	Ą	2000	0629		C.	A 19	9 <u>9</u> -2	<u>3558</u> 2	27	1999	1220		
EP	1141	021		A	ī	2001	1010		E	P 19	99-9	6835	7	1999	1220		
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	FI														
JP	2002	5362	96	T	2	2002	1029		J	P 20	00-5	89570	0	1999	1220		
PRIORITY	APP	LN.	INFO	. :					SE 1	998-	4465		Α	1998	1222		

OTHER SOURCE(S):

MARPAT 133:71091

GΙ

$$R^2$$
 R^3
 R^4
 R^4
 R^1
 R^2
 R^1
 R^2
 R^1
 R^2
 R^3

A method is disclosed for selectively enriching/removing a serum albumin from a mixt. of other compds. by contacting said mixt. with M-B-X where M is matrix, B the spacer and X the affinity ligand, with the provision that M may contain further groups X linked via a spacer. The characterizing feature is that the ligand X has been selected among serum albumin-binding structures complying with the I in which the free valence binds to the spacer B; R1-4 are selected from hydrogen, electronwithdrawing groups, such as halogens and lower alkyl groups (C1-10) that possibly are substituted with electron withdrawing groups, such as halogens; Z and Y are selected among

```
oxygen, sulfur or nitrogen, with the provision that the nitrogen may carry
a pos. charge. Also disclosed is a method for screening for ligand
structures that, when attached to an affinity matrix, selectively bind
serum albumin. The method has the characterizing feature that water-sol.
compds. that exhibit a benzene ring fused to a 5-membered heterocycle
contg. two or three heteroatoms, preferably two, selected from nitrogen,
oxygen and sulfur after having been attached to a matrix, preferably in
the 2-position, are screened for selective binding to albumin. Sepharose
4FF was activated with 1,4-bis(epoxypropoxy) butan and then coupled to
various benzimidazol-2-yl compds. and other compds. The gels were tested
for binding to human and bovine serum albumins and to human IgG.
ICM C07K014-765
9-3 (Biochemical Methods)
Section cross-reference(s): 63
serum albumin removal purifn affinity ligand;
benzimidazolyl affinity chromatog serum albumin
Adsorbents
   (affinity; removal/purifn. of serum albumins using
   matrix-immobilized affinity ligands)
Ligands
RL: BPR (Biological process); BSU (Biological study, unclassified); DEV
(Device component use); NUU (Other use, unclassified); BIOL (Biological
study); PROC (Process); USES (Uses)
   (immobilized, affinity; removal/purifn. of serum albumins
   using matrix-immobilized affinity ligands)
   (removal/purifn. of serum albumins using matrix-immobilized
   affinity ligands)
Albumins, preparation
RL: PUR (Purification or recovery); REM (Removal or disposal); PREP
(Preparation); PROC (Process)
   (serum; removal/purifn. of serum albumins using
   matrix-immobilized affinity ligands)
2425-79-8
RL: RCT (Reactant); RACT (Reactant or reagent)
   (Sepharose 4FF activation with; removal/purifn. of serum
   albumins using matrix-immobilized affinity ligands)
136109-65-4, Sepharose 4FF
RL: RCT (Reactant); RACT (Reactant or reagent)
   (epoxy activation of and reaction with ligands;
   removal/purifn. of serum albumins using matrix-immobilized
   affinity ligands)
120-53-6
          149-30-4, 2(3H)-Benzothiazolethione 583-39-1
2382-96-9, 2(3H)-Benzoxazolethione
                                     4845-58-3 5331-91-9
6325-91-3 19462-98-7 27231-36-3
37052-78-1 142313-30-2 175135-17-8
175135-18-9 175276-96-7
RL: RCT (Reactant); RACT (Reactant or reagent)
   (reaction of, with epoxy-activated Sepharose 4FF; removal/purifn. of
   serum albumins using matrix-immobilized affinity
   ligands)
136109-65-4DP, Sepharose 4FF, reaction products with ligands
RL: DEV (Device component use); NUU (Other use, unclassified); PEP
(Physical, engineering or chemical process); SPN (Synthetic preparation);
PREP (Preparation); PROC (Process); USES (Uses)
   (removal/purifn. of serum albumins using matrix-immobilized
   affinity ligands)
583-39-1 6325-91-3 19462-98-7
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27231-36-3 37052-78-1 142313-30-2

175135-17-8 175135-18-9 175276-96-7

RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, with epoxy-activated Sepharose 4FF; removal/purifn. of
serum albumins using matrix-immobilized affinity

ligands)

RN 583-39-1 HCAPLUS

CN 2H-Benzimidazole-2-thione, 1,3-dihydro- (9CI) (CA INDEX NAME)

RN 6325-91-3 HCAPLUS

CN 2H-Benzimidazole-2-thione, 1,3-dihydro-5-nitro- (9CI) (CA INDEX NAME)

RN 19462-98-7 HCAPLUS

CN 2H-Benzimidazole-2-thione, 5,6-dichloro-1,3-dihydro- (9CI) (CA INDEX NAME)

RN 27231-36-3 HCAPLUS

CN 2H-Benzimidazole-2-thione, 1,3-dihydro-5-methyl- (9CI) (CA INDEX NAME)

RN 37052-78-1 HCAPLUS

CN 2H-Benzimidazole-2-thione, 1,3-dihydro-5-methoxy- (9CI) (CA INDEX NAME)

RN 142313-30-2 HCAPLUS

CN 2H-Benzimidazole-2-thione, 5-chloro-6-fluoro-1,3-dihydro- (9CI) (CA INDEX NAME)

RN 175135-17-8 HCAPLUS

CN 2H-Benzimidazole-2-thione, 4-bromo-1,3-dihydro-6-(trifluoromethy1)- (9CI) (CA INDEX NAME)

RN 175135-18-9 HCAPLUS

CN 2H-Benzimidazole-2-thione, 4-chloro-1,3-dihydro-6-(trifluoromethyl)- (9CI) (CA INDEX NAME)

RN 175276-96-7 HCAPLUS

CN 2H-Benzimidazole-2-thione, 5-chloro-1,3-dihydro-1-(1-methylethyl)- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L21 ANSWER 3 OF 12 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 2000:12359 HCAPLUS

DOCUMENT NUMBER:

132:273792

TITLE:

Plasma protein binding of albendazole and its main

metabolite albendazole sulfoxide

AUTHOR(S):

Medina R., Liz; Garcia A., Luis; Jung C., Helgi

CORPORATE SOURCE:

Instituto Nacional de Neurologia y Neurocirugia, Fac. Quimica, UNAM, Ciudad Universitaria, DF, 04360, Mex.

SOURCE:

Revista Mexicana de Ciencias Farmaceuticas (1999),

30(3), 42-45

CODEN: RMCFDT; ISSN: 1027-3956

PUBLISHER:

Asociacion Farmaceutica Mexicana

DOCUMENT TYPE:

Journal

LANGUAGE:

Spanish

The binding of albendazole and albendazole sulfoxide to blood plasma proteins, albumin, and .alpha.1-acid glycoprotein was detd. using the equil. dialysis technique. Albendazole was bound to plasma proteins 89-92%, to albumin 80-82%, and to .alpha.1-acid glycoprotein 9-10%, whereas the binding of albendazole sulfoxide to plasma proteins was 62-67%, to albumin 33-36%, and to .alpha.1-acid glycoprotein 29-39%. binding differences may be due to lower hydrophobicity of albendazole sulfoxide than its precursor. Since the sulfoxide metabolite is responsible of the albendazole pharmacol. activity, the lower extent of its binding has no clin. significance.

CC 1-2 (Pharmacology)

IT Albumins, biological studies

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(serum; albendazole and its metabolite albendazole sulfoxide binding to blood plasma proteins in vitro)

54029-12-8, Albendazole sulfoxide 54965-21-8, IT

Albendazole

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(albendazole and its metabolite albendazole sulfoxide binding to blood plasma proteins in vitro)

IT 54029-12-8, Albendazole sulfoxide 54965-21-8,

Albendazole

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(albendazole and its metabolite albendazole sulfoxide binding to blood plasma proteins in vitro)

54029-12-8 HCAPLUS RN

Carbamic acid, [5-(propylsulfinyl)-1H-benzimidazol-2-yl]-, methyl ester CN (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 54965-21-8 HCAPLUS

Carbamic acid, [5-(propylthio)-1H-benzimidazol-2-yl]-, methyl ester (9CI) CN (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

REFERENCE COUNT:

13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L21 ANSWER 4 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1998:512080 HCAPLUS

DOCUMENT NUMBER:

130:47072

TITLE:

Sex differences in the disposition of albendazole

metabolites in sheep

AUTHOR(S):

Cristofol, Carles; Navarro, Marc; Franquelo, Carme;

Valladares, Josep-Enric; Arboix, Margarita

CORPORATE SOURCE:

Facultat de Veterinaria, Departament de Farmacologia i

de Terapeutica, UAB, Bellaterra, 08193, Spain

SOURCE:

Veterinary Parasitology (1998), 78(3), 223-231

CODEN: VPARDI; ISSN: 0304-4017

PUBLISHER:

Elsevier Science B.V.

DOCUMENT TYPE: LANGUAGE:

Journal English

Sex differences in the disposition of albendazole metabolites in sheep after oral administration of 20 mg/kg of netobimin have been studied. Some kinetic parameters of both metabolites show statistical differences between sexes; the sulfoxide and sulfone t1/2.beta. and MRT were lower in male animals than in females. Peak concns. and AUC of sulfone metabolites were higher in males suggesting a greater oxidn. rate compared with females. Urine excretion of albendazole metabolites, sulfoxide, sulfone, and amino sulfone appeared to be greater in female sheep than in males, mainly the sulfoxide metabolite. These differences between sexes can be caused by male sexual hormones, because testosterone and progesterone can induce or inhibit the microsomal Cytochrome P 450 metab. Plasma protein-binding of albendazole sulfoxide and albendazole sulfone has been studied between male and female sheep, also their binding to sheep albumin and globulins. Both albendazole metabolites readily bind to sheep albumin and globulins. Male animals show a significantly lower binding of albendazole metabolites than females. These differences could be responsible for the non-esterified fatty acids (NEFA) present in the plasma. Males have significantly higher plasma levels of NEFA than females and which may compete with for binding to albendazole metabolites.

CC 1-2 (Pharmacology)

Section cross-reference(s): 63

IT Albumins, biological studies

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(serum, binding to; sex differences in the disposition of albendazole metabolites in sheep)

IT 54965-21-8D, Albendazole, metabolites 88255-01-0, Netobimin
RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
(Biological study); PROC (Process)

(sex differences in the disposition of albendazole metabolites in sheep)

IT 54029-12-8, Albendazole sulfoxide 75184-71-3,

Albendazole sulfone 80983-34-2

RL: BPR (Biological process); BSU (Biological study, unclassified); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative); PROC (Process)

(sex differences in the disposition of albendazole metabolites in sheep)

IT 54965-21-8D, Albendazole, metabolites

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(sex differences in the disposition of albendazole metabolites in sheep)

RN 54965-21-8 HCAPLUS

CN Carbamic acid, [5-(propylthio)-1H-benzimidazol-2-yl]-, methyl ester (9CI) (CA INDEX NAME)

IT 54029-12-8, Albendazole sulfoxide 75184-71-3,

Albendazole sulfone 80983-34-2

RL: BPR (Biological process); BSU (Biological study, unclassified); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, ______nonpreparative); PROC (Process)

(sex differences in the disposition of albendazole metabolites in sheep)

RN 54029-12-8 HCAPLUS

CN Carbamic acid, [5-(propylsulfinyl)-1H-benzimidazol-2-yl]-, methyl ester (9CI) (CA INDEX NAME)

RN 75184-71-3 HCAPLUS

CN Carbamic acid, [5-(propylsulfonyl)-1H-benzimidazol-2-yl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & O & O \\ H & N & NH-C-OMe \\ \hline O & N & NH-C-OMe \\ \end{array}$$

80983-34-2 HCAPLUS RN

1H-Benzimidazol-2-amine, 5-(propylsulfonyl)- (9CI) (CA INDEX NAME) CN

$$\begin{array}{c|c}
O & H & NH2 \\
\hline
N & N & NH2
\end{array}$$

REFERENCE COUNT:

22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L21 ANSWER 5 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1997:344326 HCAPLUS

DOCUMENT NUMBER:

127:26008

TITLE:

Negatively charging electrostatographic toner

containing 2-substituted imidazole derivative charge ____ __ ____

controller

INVENTOR(S):

Takahashi, Toshihiko; Tanaka, Katsuhiko; Nagatsuka,

APPLICATION NO. DATE

Takayuki

PATENT ASSIGNEE(S):

Canon K. K., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 12 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

KIND DATE

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT NO.

PATENT INFORMATION:

	JP 09080819	A2	19970328	JP 1995-257217	19950911		
	JP 3382428	B2	20030304				
PRIO	RITY APPLN. INFO .:			JP 1995-257217	19950911		
OTHE	R SOURCE(S):	MA	RPAT 127:260	800			
AB	The toner contain	s an	imidazole co	ompd. having an elec	tron-withdrawing		
substituent at the 2nd position. The toner showed rapid and enough							
	charging and long	shel	f life.		· .		
IC	ICM G03G009-097						
	ICS G03G009-08						
CC	74-3 (Radiation C	hemis	try, Photoch	nemistry, and Photog	raphic and Other		
	Reprographic Proc	esses)	-	_		
ST	neg charging elec	trost	atog toner i	midazole; electron			
withdrawing substituent imidazole electrophotog toner; charge							
	controller imidaz	ole e	lectrostato	g toner	-		
IT	50832-48-9 8176	9-47-	3 131769-26-	-1 189338-47-4			
	RL: TEM (Technica	l or	engi.neered m	naterial use); USES	(Uses)		

(neg.-charging electrostatog. toner contg. imidazole deriv. charge controller showing rapid and enough charging and long shelf life)

IT 131769-26-1 189338-47-4

RL: TEM (Technical or engineered material use); USES (Uses)

(neg.-charging electrostatog. toner contg. imidazole deriv. charge controller showing rapid and enough charging and long shelf life)

RN 131769-26-1 HCAPLUS

CN 1H-Benzimidazole, 2-(pentadecafluoroheptyl)- (9CI) (CA INDEX NAME)

RN 189338-47-4 HCAPLUS

CN 1H-Benzimidazole, 5-methyl-2-(pentadecafluoroheptyl)- (9CI) (CA INDEX NAME)

L21 ANSWER 6 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1995:554835 HCAPLUS

DOCUMENT NUMBER:

123:3079

TITLE:

Charge transfer chromatographic study of the **binding** of commercial pesticides to various

albumins

AUTHOR(S):

Cserhati, Tibor; Forgacs, Esther

CORPORATE SOURCE:

Central Research Institute for Chemistry, Hungarian Academy of Sciences, P.O. Box 17, Budapest, 1525,

Hung.

SOURCE:

Journal of Chromatography, A (1995), 699(1 + 2),

285-90

CODEN: JCRAEY; ISSN: 0021-9673

PUBLISHER:

Elsevier Journal

DOCUMENT TYPE: LANGUAGE:

Journal English

The interaction of 28 com. pesticides with human and bovine serum albumin as well as with egg albumin was studied by charge-transfer reversed-phase thin-layer chromatog, and the relative strength of the interaction was calcd. Only one pesticide interacted with egg albumin whereas the majority of pesticides bound both to bovine and human serum albumins. Stepwise regression anal. proved that the hydrophobicity parameters of pesticides exert a significant impact on their capacity to bind to serum albumins. These findings support the hypothesis that the binding of pesticides to albumins may involve hydrophilic forces occurring between the corresponding apolar substructures of pesticides and amino acid side chains. No linear correlation was found between the capacities of human and bovine serum albumins to bind pesticides.

CC 4-4 (Toxicology)

Section cross-reference(s): 5

ST pesticide binding albumin charge transfer chromatog

IT Pesticides

(charge transfer chromatog. study of **binding** of com. pesticides to various **albumins**)

IT Albumins, biological studies

Ovalbumins

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(charge transfer chromatog. study of binding of com.

pesticides to various albumins)

IT Chromatography, column and liquid

(charge-transfer, charge transfer chromatog. study of binding of com. pesticides to various albumins)

115-29-7, Endosulfan IT 80-33-1, Chlorfenson 330-55-2, Linuron 886-50-0, Terbutryn 957-51-7, Diphenamid 1912-24-9, Atrazin 2032-65-7, Methiocarb 2164-08-1, Lenacil 2425-06-1, Captafol 3878-19-1, Fuberidazole 4658-28-0, Aziprotryne 5234-68-4, Carboxin 5902-51-2, Terbacil 5915-41-3, Terbutylazine 13360-45-7, Chlorbromuron 15545-48-9, Chlorotoluron **17804-35-2**, Benomyl 23564-05-8, Thiophanate-methyl 26225-79-6, Ethofumesate 34123-59-6, Isoproturon 57966-95-7 67747-09-5, Prochloraz 69327-76-0, Buprofezin 74115-24-5, 74782-23-3, Oxabetrinil Clofentezine 76674-21-0, Flutriafol 82097-50-5, Triasulfuron 77732-09-3, Oxadixyl RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(charge transfer chromatog. study of binding of com.

pesticides to various albumins)

IT 17804-35-2, Benomyl

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(charge transfer chromatog. study of binding of com.

pesticides to various albumins)

RN 17804-35-2 HCAPLUS

CN Carbamic acid, [1-[(butylamino)carbonyl]-1H-benzimidazol-2-yl]-, methyl ester (9CI) (CA INDEX NAME)

L21 ANSWER 7 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1994:425557 HCAPLUS

DOCUMENT NUMBER:

121:25557

TITLE:

Copper(II) complexes of novel tripodal **ligands** containing phenolate and benzimidazole/pyridine pendants: synthesis, structure, spectra and

electrochemical behavior

AUTHOR(S):

Uma, Rajendran; Viswanathan, Rathinam; Palaniandavar,

Mallayan; Lakshminarayanan, M.

CORPORATE SOURCE:

Dep. Chem., Bharathidasan Univ., Tiruchirapalli, 620

024, India

Journal of the Chemical Society, Dalton Transactions: SOURCE:

Inorganic Chemistry (1972-1999) (1994), (8), 1219-26

CODEN: JCDTBI; ISSN: 0300-9246

DOCUMENT TYPE: Journal LANGUAGE: English

Mononuclear Cu(II) complexes of tri- and tetra-dentate tripodal ligands 2-HO-5-NO2C6H3NRCH2R1 (R = H, R1 = 2-benzimidazolyl (R2), 2-pyridyl (R3); R = R1 = R2, R3), 2-H0-5-N02C6H3CH2NHCH2CH2R2, (2-H0-5-N02C6H3CH2)2NR (R = R2, R3) and HO-5-NO2C6H3CH2NHR3 isolated. They are [CuL(X)].cntdot.nH2O, [CuL(H2O)]X.cntdot.nH2O or [CuL].cntdot.nH2O where X = Cl-, ClO4-, N3- or NCS- and n = 0-4. The electronic spectra of all the complexes exhibit a broad absorption band around 14,000 cm-1 and the polycryst. as well as the frozen-soln. EPR spectra are axial, indicating square-based geometries. The crystal structure of [CuLCl] [HL = (2-hydroxy-5-nitrobenzyl)bis(2pyridylmethyl)amine] revealed a square-pyramidal geometry around CuII. The mononuclear complex crystallizes in the triclinic space group P.hivin.1 with a 6.938(1), b 11.782(6), c 12.678(3) .ANG. and .alpha. 114.56(3), .beta. 92.70(2), .gamma. 95.36(2).degree.. The coordination plane is comprised of 1 tertiary amine and 2 pyridine nitrogens and a chloride ion. The phenolate ion unusually occupies the axial site, possibly due to the electron-withdrawing p-nitro group. The enhanced .pi. delocalization involving the p-nitrophenolate donor elevates the E1/2 values. The spectral and electrochem. results suggest the order of donor strength as nitrophenolate < pyridine < benzimidazole in the tridentate and nitrophenolate < benzimidazole < pyridine in the tetradentate ligand complexes.

78-7 (Inorganic Chemicals and Reactions)

Section cross-reference(s): 72, 75

4499-07-4, 2-(2-Aminoethyl)benzimidazole dihydrochloride 5993-91-9, 2-Aminomethylbenzimidazole dihydrochloride RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with chloromethylnitrophenol)

IT 4499-07-4, 2-(2-Aminoethyl)benzimidazole dihydrochloride 5993-91-9, 2-Aminomethylbenzimidazole dihydrochloride

RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with chloromethylnitrophenol)

4499-07-4 HCAPLUS RN

1H-Benzimidazole-2-ethanamine, dihydrochloride (9CI) (CA INDEX NAME) CN

$$\begin{array}{c|c} H & \operatorname{CH_2-CH_2-NH_2} \\ \hline & N & \end{array}$$

O2 HC1

RN5993-91-9 HCAPLUS

1H-Benzimidazole-2-methanamine, dihydrochloride (9CI) (CA INDEX NAME) CN

$$\underbrace{ \overset{\text{H}}{\underset{\text{N}}{\bigvee}} \text{CH}_2 - \text{NH}_2 }_{\text{H}}$$

●2 HCl

L21 ANSWER 8 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1992:58905 HCAPLUS

DOCUMENT NUMBER:

116:58905

TITLE:

Mono- and bis(2-nitroguanidino)benzenes and some of

their amino and nitro derivatives

AUTHOR(S):

Luk'yanov, O. A.; Mel'nikova, T. G.; Shagaeva, M. E.

CORPORATE SOURCE:

Inst. Org. Khim. im. Zelinskogo, Moscow, USSR

SOURCE:

Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya

(1991), (11), 2581-7

CODEN: IASKA6; ISSN: 0002-3353

DOCUMENT TYPE:

Journal

LANGUAGE:

Russian

OTHER SOURCE(S):

CASREACT 116:58905

GΙ

- AΒ Reaction of MeSC(NH2):NNO2 (I) with arylamines ArNH2 (Ar = Ph, 3- and 4-H2NC6H4) at 60-80.degree. afforded the corresponding mono(nitroguanidino) derivs. ArNHC(NH2):NNO2 in 76, 89, and 80% yields, resp. Reaction of I with o-phenylenediamine afforded (nitramino)benzimidazole II, derived from the corresponding primary product ArNHC(NH2):NNO2 (Ar = 2-H2NC6H4, III) under the reaction conditions. III itself was synthesized at lower temp. in the reaction of o-phenylenediamine with 1-methyl-1-nitroso-2-nitroguanidine, and was converted in 93% yield to II at 150-160.degree.. Bis(nitroguanidino) substitution in ArNH2 was accomplished at higher temp. and for longer reaction duration, testifying to the deactivating effect of the electron-accepting nitroguanidino group on the reaction of the remaining nitro group. Alternative synthetic routes for (nitroaryl)-2nitroguanidines involved oxidn. of the corresponding (aminoaryl) and nitration of the corresponding aryl derivs.
- CC 25-21 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds) Section cross-reference(s): 28
- IT Nitration

(of (nitroguanidino) benzenes contq. deactivating electronwithdrawing groups)

ITRegiochemistry

(of nitration of (nitroguanidino)benzenes contg. deactivating
electron-withdrawing groups)

IT 138416-36-1P 138416-41-8P 138416-42-9P 138416-45-2P

138416-46-3P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

IT 138416-36-1P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

RN 138416-36-1 HCAPLUS

CN 1H-Benzimidazol-2-amine, N-nitro- (9CI) (CA INDEX NAME)

L21 ANSWER 9 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1988:17530 HCAPLUS

DOCUMENT NUMBER:

108:17530

TITLE:

Prevention by thioethers of the hepatotoxicity and

covalent binding to macromolecules of

N-hydroxy-2-acetylaminofluorene and its sulfate ester

in rat liver in vivo and in vitro

AUTHOR(S):

Van den Goorbergh, J. A. M.; De Wit, H.; Tijdens, R.

B.; Mulder, G. J.; Meerman, J. H. N.

CORPORATE SOURCE:

Sylvius Lab., Univ. Leiden, Leiden, 2300 RA, Neth.

SOURCE:

Carcinogenesis (1987), 8(2), 275-9

DOCUMENT TYPE:

CODEN: CRNGDP; ISSN: 0143-3334 Journal

LANGUAGE:

English

Ι

GT

To find potentially effective compds. that could prevent the covalent binding of the carcinogen N-hydroxy-2-acetylaminofluorene (N-OH-AAF) (I) to rat liver macromols. in vivo, the prevention of the covalent binding to RNA of the sulfate ester of N-OH-AAF by a series of thioethers was investigated in vitro. The most effective thioethers, which inhibited the covalent binding by .gtoreq.70% were studied for their protection against acute hepatotoxicity of N-OH-AAF in the rat in vivo. Three of these thioethers, thiazolidine, Me 4-(methylthio)benzoate, and 2-(methylthio)benzimidazole, significantly decreased the hepatotoxicity of N-OH-AAF by 45, 71, and 83%, resp. The effects of these thioethers on the covalent binding of N-OH-AAF to cellular macromols. in vivo were also

studied. Me 4-(methylthio)benzoate and 2-(methylthio)benzimidazole decreased the adduct formation of N-OH-AAF to DNA by 54 and 44%, resp., but had no effect on protein adduct formation. Only 2-(methylthio)benzimidazole caused a slight decrease (23%) in the AAF-protein adduct formation. AAF and Me 4-(methylsulfinyl)benzoate were the main products in the incubation of Me 4-(methylthio)benzoate with AAF-N-sulfate in vitro. This suggests that the thioether attacks the nitrenium ion which is formed by spontaneous breakdown of AAF-N-sulfate; the formation of a sulfonium-AAF conjugate is postulated which decomps. into AAF and a sulfinyl compd.

CC 4-6 (Toxicology)

IT Albumins, biological studies RL: BIOL (Biological study)

(hydroxyacetylaminofluorene sulfate covalent binding to,

thioethers effect on)

147-84-2, Diethyldithiocarbamic acid, biological studies 444-27-9, ΙT Thiazolidine 4-carboxylic acid 504-78-9, Thiazolidine 3795-79-7, Methyl 4-(methylthio)benzoate 7152-24-1, 2-

(Methylthio) benzimidazole

RL: BIOL (Biological study)

(hydroxyacetylaminofluorene toxicity to liver response to, covalent binding of hydroxyacetylaminofluorene sulfate to RNA in relation to)

IT 7152-24-1, 2-(Methylthio)benzimidazole

RL: BIOL (Biological study)

(hydroxyacetylaminofluorene toxicity to liver response to, covalent binding of hydroxyacetylaminofluorene sulfate to RNA in relation to)

7152-24-1 HCAPLUS RN

1H-Benzimidazole, 2-(methylthio)- (9CI) (CA INDEX NAME) CN

L21 ANSWER 10 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

CORPORATE SOURCE:

1984:608793 HCAPLUS

DOCUMENT NUMBER:

101:208793

TITLE:

Monoclonal antibodies specific for .beta.-adrenergic

AUTHOR(S):

Chamat, Soulaima; Hoebeke, Johan; Strosberg, A. Donny

Lab. Mol. Immunol., Inst. Jacques Monod, Paris,

F-75251, Fr.

SOURCE:

Journal of Immunology (1984), 133(3), 1547-52

CODEN: JOIMA3; ISSN: 0022-1767

DOCUMENT TYPE:

Journal

LANGUAGE:

English

After somatic cell fusion between splenocytes of immunized BALB/c mice and NS-1 myeloma cells, 8 clones were obtained secreting anti-alprenolol antibodies as characterized by means of an ELISA. Four of these were subcloned and were studied further. The assocn. const. for alprenolol ranged from 1.9 .times. 106 M-1 to 24 .times. 106 M-1. Competitive inhibition of [3H]-1-dihydroalprenolol binding revealed cross-reactivity with .beta.-adrenergic ligands, with a higher avidity for antagonists than for agonists. Two of the antibodies had a higher affinity for the 1-isomer than for the d-isomer. The most stereospecific of these

antibodies showed only affinity for .beta.-adrenergic antagonists and for the agonist isoproterenol. The other recognized both .beta.-adrenergic antagonists and agonists; it also showed an increase in tryptophan fluorescence after ligand binding. This property was used for the physicochem. study of the hapten-antibody interaction.

CC 15-3 (Immunochemistry)

IT Antibodies

RL: BIOL (Biological study)

(monoclonal, to alprenolol, .beta.-adrenergic ligand

specificity of)

IT 51-31-0 51-41-2 51-43-4 2964-04-7 4199-09-1 5051-22-9 6673-35-4 18559-94-9 60106-89-0 72332-33-3 **81047-99-6**

RL: BIOL (Biological study)

(alprenolol-specific monoclonal antibody binding to)

IT 13655-52-2D, albumin conjugates

RL: BIOL (Biological study)

(monoclonal antibodies to, .beta.-adrenergic ligand

specificity of)

IT 81047-99-6

RL: BIOL (Biological study)

(alprenolol-specific monoclonal antibody binding to)

RN 81047-99-6 HCAPLUS

CN 2H-Benzimidazol-2-one, 4-[3-[(1,1-dimethylethyl)amino]-2-hydroxypropoxy]-1,3-dihydro-(9CI) (CA INDEX NAME)

L21 ANSWER 11 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1977:116700 HCAPLUS

DOCUMENT NUMBER: 86:116700

TITLE: Acyl migrations in diacyl derivatives of

2-methylmercaptobenzimidazole. A model of biotin

AUTHOR(S):

Ohno, A.; Morishita, T.; Oka, S.

CORPORATE SOURCE: SOURCE:

Inst. Chem. Res., Kyoto Univ., Uji, Japan
Bioorganic Chemistry (1976), 5(4), 383-91

CODEN: BOCMBM; ISSN: 0045-2068

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB Diacyl derivs. of 2-methylmercaptobenzimidazole undergo tautomerization. Thermodn. predominancy of 1 isomer over the others depends on the substituents on carbonyl groups. Electron-withdrawing and electron-releasing substituents favor different isomeric configurations. The migration was extended to include the carboethoxy group and the results are discussed in relation to the mechanism of biotin-dependent enzymic carboxylation.

CC 7-4 (Enzymes)

IT 7152-24-1D, diacyl derivs.

RL: BIOL (Biological study)

(acyl migrations in, electron-releasing and electron

-withdrawing substituents in relation to) IT62312-50-9 RL: BIOL (Biological study) (ethoxycarbonyl of, migration of) IT 5268-66-6 RL: FORM (Formation, nonpreparative) (formation of, by acyl migration from ethoxycarbonylacetonylthiobenzimi dazole) IT **5268-65-5P 16458-79-0P** 18606-28-5P 51949-53-2P 52026-33-2P 62312-51-0P 62312-52-1P 62312-53-2P 62312-54-3P RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of) 21547-79-5 IT 5429-62-9 RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with acetyl chloride) IT 5268-67-7 RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with benzoyl chloride) IT 7152-24-1D, diacyl derivs. RL: BIOL (Biological study) (acyl migrations in, electron-releasing and electron -withdrawing substituents in relation to) 7152-24-1 HCAPLUS RN1H-Benzimidazole, 2-(methylthio)- (9CI) (CA INDEX NAME) CN

$$\begin{array}{c|c} & \circ & \circ \\ \parallel & \circ & \circ \\ & N & \circ & \\ & N & \circ & \circ \\ & N & \circ & \\ & N & \circ & \circ \\ & N & \circ & \\ & N & \circ & \circ \\ & N & \circ & \\ & N & \circ & \circ \\ & N & \circ$$

IT 5268-66-6
 RL: FORM (Formation, nonpreparative)
 (formation of, by acyl migration from ethoxycarbonylacetonylthiobenzimi dazole)
RN 5268-66-6 HCAPLUS
CN Butanoic acid, 2-(1H-benzimidazol-2-ylthio)-3-oxo-, ethyl ester (9CI) (CA INDEX NAME)

IT 5268-65-5P 16458-79-0P 62312-54-3P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

RN 5268-65-5 HCAPLUS

CN 2,4-Pentanedione, 3-(1H-benzimidazol-2-ylthio)- (9CI) (CA INDEX NAME)

RN 16458-79-0 HCAPLUS

CN 1H-Benzimidazole, 1-acetyl-2-[(2-oxopropyl)thio]- (9CI) (CA INDEX NAME)

RN 62312-54-3 HCAPLUS

CN Acetic acid, [(1-acetyl-1H-benzimidazol-2-yl)thio]-, ethyl ester (9CI) (CA INDEX NAME)

IT 5429-62-9

RL: RCT (Reactant); RACT (Reactant or reagent)

(reaction of, with acetyl chloride)

RN 5429-62-9 HCAPLUS

CN Acetic acid, (1H-benzimidazol-2-ylthio)-, ethyl ester (9CI) (CA INDEX NAME)

IT 5268-67-7

RL: RCT (Reactant); RACT (Reactant or reagent)
 (reaction of, with benzoyl chloride)

RN 5268-67-7 HCAPLUS

CN 2-Propanone, 1-(1H-benzimidazol-2-ylthio)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} H & S-CH_2-C-Me \\ \hline \\ N & N \end{array}$$

L21 ANSWER 12 OF 12 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1973:522311 HCAPLUS

DOCUMENT NUMBER:

79:122311

TITLE:

Effect of some uncoupling agents, ionophorous agents,

and inhibitors on the fluorescence of ANS

[1-anilino-8-naphthalenesulfonate] bound to bovine

serum albumin

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The uncouplers, tetrachlorotrifluoromethyl benzimidazole (TTFB) [
2338-29-6], carbonyl cyanide p-trifluoromethaoxyphenyl hydrazone
(I) [370-86-5] and carbonyl cyanide m-chlorophenyl hydrazone (CCCP)
[555-60-2] considerably decreased the fluorescence of 1-anilino-8naphthalenesulfonate (ANS) [82-76-8] bound to bovine serum albumin. TTFB
exhibited satn., whereas I and CCCP eliminated all the bovine serum
albumin enhancement of ANS fluorescence. Ionophorous agents, such as
nigericin [28380-24-7], and the ATPase inhibitor, oligomycin [1404-19-9],
increased fluorescence. The interaction of bovine serum albumin with the
uncouplers appears to affect the ANS binding site and to decrease the amt.
of probe bound.

CC 3-13 (Biochemical Interactions)

serum albumin ANS binding uncoupler; ionophorous agent albumin ANS binding; anilinonaphthalenesulfonate binding albumin

RL: PRP (Properties)

(albumin-anilinonaphthalenesulfonate complex fluorescence response to)

IT 2338-29-6

RL: PRP (Properties)

(albumin-anilinonaphthalenesulfonate complex fluorescence response to)

RN 2338-29-6 HCAPLUS

CN 1H-Benzimidazole, 4,5,6,7-tetrachloro-2-(trifluoromethyl)- (9CI) (CA INDEX NAME)